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TRANSMITTAL LETTER TO THE UNITED STATES ATTORNEY'S DOCKET NUMBER_48985 DESIGNATED/ELECTED OFFICE (DO/EO/US) 674047 CONCERNING A FILING UNDER 35 U.S.C. 371 U.S. APPLICATION NO. (If known, INTERNATIONAL APPLICATION NO. INTERNATIONAL FILING DATE PRIORITY DATE CLAIMED PCT/EP 99/02544 April 15, 1999 April 27, 1998 TITLE OF INVENTION: METHOD FOR THE CATALYTIC DECOMPOSITION OF N₂O APPLICANT(S) FOR DO/EO/US Volker SCHUMACHER, Gert BÜRGER, Thomas FETZER, Michael BAIER, Michael HESSE Applicant herewith submits to the United States Designated/Elected Office (DO/EO/US) the following items and other information: 1. /X/ This is a FIRST submission of items concerning a filing under 35 U.S.C. 371. 2.11 This is a SECOND or SUBSEQUENT submission of items concerning a filing under 35 U.S.C. 371. This express request to begin national examination procedures (35 U.S.C.371(f)) at any time rather than delay examination 3./X/ until the expiration of the applicable time limit set in 35 U.S.C. 371(b) and PCT Articles 22 and 39(1). 4.// A proper Demand for International Preliminary Examination was made by the 19th month from the earliest claimed priority date A copy of the International Application as filed (35 U.S.C. 371(c)(2)). 5. /X/ a./X/ is transmitted herewith (required only if not transmitted by the International Bureau). h// has been transmitted by the International Bureau C./ / is not required, as the application was filed in the United States Receiving Office (RO/USO). 6. /X/ A translation of the International Application into English (35 U.S.C. 371(c)(2)). 7. // Amendments to the claims of the International Application under PCT Article 19 (35 U.S.C. 371(c)(3)). a.// are transmitted herewith (required only if not transmitted by the International Bureau). b.// have been transmitted by the International Bureau. have not been made; however, the time limit for making such amendments has NOT expired. c// d.// have not been made and will not be made. A translation of the amendments to the claims under PCT Article 19(35 U.S.C. 371(c)(3)), 8.//

/X/ An oath or declaration of the inventor(s)(35 U.S.C. 171(c)(4)).

10.// A translation of the annexes to the International Preliminary Examination Report under PCT Article 36 (35 U.S.C. 371(c)(5)). Items 11. to 16. below concern other document(s) or information included:

11.// An Information Disclosure Statement under 37 CFR 1.97 and 1.98.

12./X/ An assignment document for recording. A separate cover sheet in compliance with 37 CFR 3.28 and 3.31 is included.

13./X/ A FIRST preliminary amendment.

// A SECOND or SUBSEQUENT preliminary amendment.

14.// A substitute specification.

15.// A change of power of attorney and/or address letter.

16./X/ Other items or information.
International Search Report
International Preliminary Examination Report

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SEND ALL CORRESPONDENCE TO: KEIL & WEINKAUF 1101 Connecticut Ave., N.W.

Washington, D. C. 20036

SIGNATURE

Herbert B. Keil NAME 18,967 Registration No. IN THE UNITED STATES PATENT AND TRADEMARK OFFICE
In re the U.S. Nationalization Application
of PCT/EP99/02544 of)

SCHUMACHER et al.)

Intl. Filing Date: April 15, 1999)

US Serial No.: TO BE ASSIGNED)

Filed: Herewith)

For: METHOD OF CATALYTIC DECOMPOSITION OF N2O

Honorable Commissioner of Patents and Trademarks Washington, D.C. 20231

PRELIMINARY AMENDMENT

Sir:

Prior to examination of the above-identified U.S. National Stage application, kindly amend the application as follows.

IN THE CLAIMS

Please cancel claim 1-10.

Please add claims 11-18 as follows

- 11. A reactor for the catalytic oxidation of ammonia to nitrogen oxides, which contains a noble metal gauze catalyst and a heat exchanger in that order in the direction of flow and has a catalyst for the decomposition of N₂O which can be prepared by combining CuAl₂O₄ with tin, lead and/or an element of main group II or transition group II of the Periodic Table of the Elements as oxide or salt or in elemental form and subsequently calcining the mixture at from 300 to 1300°C and a pressure in the range from 0.1 to 200 bar located between the noble metal gauze catalyst and the heat exchanger.
- 12. A reactor as claimed in claim 11, wherein a noble metal recovery gauze is located between the noble metal gauze catalyst and the catalyst for the decomposition of N_2O .

- 13. A reactor as claimed in claim 11, wherein the catalyst for the decomposition of N_2O is installed as a fixed bed having a height of from 2 to 50 cm.
 - 14. An apparatus for preparing nitric acid from ammonia, comprising in this order
 - a reactor as claimed in claim 11,
 - an absorption unit for the absorption of nitrogen oxides in an aqueous medium and, if desired,
 - a reduction unit for the selective catalytic reduction of nitrogen oxides.
 - 15. An apparatus for preparing nitric acid from ammonia, comprising in this order
 - a reactor as claimed in claim 12,
 - an absorption unit for the absorption of nitrogen oxides in an aqueous medium and, if desired,
 - a reduction unit for the selective catalytic reduction of nitrogen oxides.
- 16. A process for the catalytic decomposition of N₂O in a gas mixture obtained in the preparation of nitric acid by catalytic oxidation of ammonia in a reactor having a noble metal gauze catalyst and a heat exchanger in that order in the flow direction, where N₂O is decomposed catalytically over a catalyst for the decomposition of N₂O located between the noble metal catalyst and the heat exchanger so that the hot gas mixture obtained from the catalytic oxidation of ammonia is brought into contact with the catalyst for the decomposition of N₂O prior to subsequent cooling, wherein the catalyst for the decomposition of N₂O can be prepared by combining CuAl₂O₄ with tin, lead or an element of main group II or transition group II of the Periodic Table of Elements as oxide or salt or in elemental form and subsequently calcining the mixture at from 300 to 1300°C and a pressure in the range from 0.1 to 200 bar.
- 17. A process as claimed in claim 16, wherein the residence time over the catalyst for the decomposition of N₂O is less than 0.1 s.

18. A process as claimed in claim 16, wherein the decomposition of N_2O is carried out at from 600 to 950°C and/or at a pressure in the range from 1 to 15 bar.

REMARKS

The claims have been amended to eliminate multiple dependency and to place them in better form for U.S. practice. No new matter has been added. Favorable action on the application is solicited.

Please charge any shortage in fees due in connection with the filing of this paper, to Deposit Account No. 11-0345. Please credit any excess fees to such deposit account.

Respectfully submitted,

KEIL & WEINKAUF

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AS ORIGINALLY FILED

Catalytic decomposition of N₂O

The present invention relates to a process for the catalytic decomposition of N_2O in a gas mixture obtained in the preparation of nitric acid by catalytic oxidation of ammonia. The invention further relates to a reactor which is suitable for carrying out the process.

In the industrial preparation of nitric acid by the Ostwald process, ammonia is reacted with oxygen over a noble metal catalyst to form oxides of nitrogen which are subsequently absorbed in water. In this process, ammonia and oxygen or air are reacted at from 800 to 955°C over a catalyst gauze comprising noble metals in a reactor. The catalyst gauze generally comprises platinum and rhodium as active metals. In the catalytic reaction, ammonia is firstly oxidized to nitrogen monoxide which is subsequently further oxidized by oxygen to give nitrogen dioxide or dinitrogen tetroxide. The gas mixture obtained is cooled and then passed to an absorption tower in which nitrogen dioxide is absorbed in water and converted into nitric acid. The reactor for the catalytic combustion of ammonia also contains, downstream of the catalyst gauze, a recovery gauze for depositing and thus recovering catalyst metals which have been vaporized at the high reaction temperatures. A heat exchanger is located downstream of the recovery gauze to cool the gas mixture obtained. Absorption is carried out outside the actual reactor in a separate absorption column.

The combustion and the absorption can be carried out at the same pressure level. It is possible to employ an intermediate pressure of from about 230 to 600 kPa or a high pressure of from about 700 to 1100 kPa. In the case of a process with two pressure stages, the absorption is carried out at a higher pressure than the combustion. The pressure in the combustion is then from about 400 to 600 kPa and the pressure in the absorption is from about 900 to 1400 kPa.

An overview of the Ostwald process may be found, for example, in Ullmann's Encyclopedia of Industrial Chemistry, 5th edition, Volume A 17, pages 293 to 339 (1991).

5 The combustion of ammonia forms not only nitrogen monoxide and nitrogen dioxide or dinitrogen tetroxide but generally also N₂O (dinitrogen monoxide) as by-product. In contrast to the other oxides of nitrogen formed, N₂O is not absorbed by the water during the absorption step. If no further step for removing N₂O is provided, N₂O can be emitted into the environment in a concentration of from about 500 to 3000 ppm in the waste gas.

Since N₂O is a greenhouse gas and participates in the depletion of the ozone layer, very substantial removal from the waste gas is desirable. A number of methods of removing N₂O from waste gas streams have been described.

DE-A-195 33 715 describes a process for removing nitrogen oxides from a gas stream, in which the nitrogen oxides apart from N_2O are absorbed in an absorption medium and remaining N_2O is subsequently decomposed catalytically at from 700 to $800^{\circ}C$ in a decomposition reactor. Since nitrogen oxides can be formed in this decomposition, a selective catalytic reduction (SCR) can follow.

US 5,478,549 describes a process for preparing nitric acid by the Ostwald method, in which the N₂O content is reduced by passing the gas stream after the oxidation over a catalyst bed of zirconium oxide at a temperature of at least 600°C. However, the precise position of the catalyst bed is not indicated. It is only stated that zirconium oxide in the form of cylindrical pellets is located below the recovery gauze. The space velocity in the reactor is 30,000 h at 4 bar and 600°C.

EP-B 0 359 286 describes a process for the reduction of N_2O . For this purpose, a reactor for carrying out the Ostwald process is modified in such a way that the gases obtained after the catalytic combustion are subjected to a retention time of from 0.1 to 3 seconds before cooling by means of the heat exchanger. If desired, a catalyst for the selective decomposition of N_2O can be additionally provided.

35 It is an object of the present invention to provide a process and an apparatus for preparing nitric acid by the Ostwald method, by means of which the N₂O content of the waste gases can be reduced very effectively and at low cost. The process

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should also be able to be integrated into existing plants and preferably increase the proportion of useful product. In addition, additional heating and cooling steps for the waste gas should be avoided, so that the removal of N_2O can be carried out economically.

We have found that this object is achieved by a reactor for the catalytic oxidation of ammonia to nitrogen oxides, which contains a noble metal gauze catalyst and a heat exchanger in that order in the direction of flow and has a catalyst for the decomposition of N_2O located between the noble metal gauze catalyst and the heat exchanger.

The object is also achieved by a process for the catalytic decomposition of N_2O in a gas mixture obtained in the preparation of nitric acid by catalytic oxidation of ammonia, where the N_2O is decomposed catalytically over a catalyst for the decomposition of N_2O , wherein the hot gas mixture obtained from the catalytic oxidation of ammonia is brought into contact with the catalyst for the decomposition of N_2O prior to subsequent cooling.

According to the present invention, it has been found that N₂O can be reacted directly in the reactor for the catalytic oxidation of ammonia when a suitable catalyst is located between the noble metal gauze catalyst and the heat exchanger. In this way, N₂O formed as by-product is decomposed immediately after it is formed. The decomposition occurs at the temperature prevailing in the catalytic oxidation of ammonia. Heating or cooling of the gaseous reaction mixture is thus unnecessary. The catalyst for the decomposition of N₂O which is used according to the present invention is located directly in the reactor, preferably between the position of a noble metal recovery gauze located downstream of the noble metal catalyst and the position of the heat exchanger. Reactors for the Ostwald process are usually provided with inserts for accommodating the noble metal catalyst and the noble metal recovery gauze. These reactors can easily be modified by additionally providing a holder for the N₂O decomposition catalyst.

The low catalyst bed height required according to the present invention allows installation in existing reactors without great rebuilding of the reactors. Thus, existing reactors can be modified to enable the process of the present invention to be carried out, without replacement of the reactor being necessary. The Ostwald process can be carried out at one pressure level or at two pressure levels, as

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described above. The height of the catalyst bed is preferably from 2 to 50 cm, particularly preferably from 5 to 10 cm. In production, the residence time over the catalyst is preferably less than 0.1 s. The pressure drop caused by installation of the catalyst is therefore very low, a small amount of catalyst can be employed, and the gas has to be held at a high temperature level for only a short time after the oxidation, so that secondary reactions can largely be suppressed.

According to the present invention, the decomposition of N₂O is carried out in the reactor for the oxidation of ammonia at the oxidation temperature, generally at a temperature in the range from 600 to 950°C, preferably from 800 to 930°C, in particular from 850 to 920°C. The pressure is, depending on the pressure level at which the Ostwald process is carried out, generally from 1 to 15 bar.

As noble metal gauze catalyst, it is possible to use any noble metal gauze catalyst suitable for the catalytic oxidation of ammonia. The catalyst preferably comprises platinum and possibly rhodium and/or palladium as catalytically active metals.

The noble metal recovery gauze is preferably made of palladium. The catalyst used according to the present invention for the decomposition of N₂O is preferably selected from among catalysts which still have sufficient activity at above 900°C to decompose N₂O at this temperature in the presence of NO and/or NO₂. Catalysts which are suitable for the purposes of the present invention are, for example, binary oxides such as MgO, NiO, ZnO, Cr₂O₃, TiO₂, WO_x, SrO, CuO/Cu₂O, Al₂O₃, Se₂O₃, MnO₂ or V₂O₅, if desired doped with metal oxides, lanthanide complexes such as La₂NiO₄, La₂CuO₄, Nd₂CuO₄ and multinary oxide compounds thereof, spinels, ternary perovskites, and also oxidic systems such as CuO-ZuO-Al₂O₃, CoO-MgO, CoO-La₂O₃, CO-ZuO, NiO-MoO₃ or metals such as Ni, Pd, Pt, Cu, Ag, Preference is given to using a catalyst as described in DE-A-43 01 470.

30 Such a catalyst can be prepared, for example, by combining CuAl₂O₄ with tin, lead and/or an element of main group II or transition group II of the Periodic Table of the Elements as oxide or salt or in elemental form, and subsequently calcining the mixture at from 300 to 1300°C and a pressure in the range from 0.1 to 200 bar. The catalyst can have any suitable shape. It is preferably used in extrudate form, in particular in the form of star extrudates. The preferred diameter of the extrudates is from 2 to 10 mm, particularly preferably from 3 to 6 mm. The catalyst can also be used in other forms, particularly also in the form of a honeycomb catalyst.

The catalyst is preferably prepared using zinc, magnesium, calcium, strontium and/or barium as oxide or salt or in elemental form in addition to CuAl₂O₄. The catalyst is preferably free of noble metals.

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To prepare the catalyst, use is made of CuAl₂O₄ of which from 1 to 100% by weight, preferably from 10 to 100% by weight, particularly preferably from 80 to 100% by weight, is present as spinel. It is particularly preferably completely in the form of spinel. Mixing with tin, lead and/or an element of main group II or transition group II of the Periodic Table of the Elements is preferably carried out at from 500 to 1200°C, particularly preferably from 600 to 1100°C, and preferably at pressures of from 0.5 to 10 bar, particularly preferably at atmospheric pressure. Mixing can be carried out, for example, by spraying, mechanical mixing, stirring or kneading of the milled solid of the composition CuAl₂O₄. Particular preference is given to impregnation of the unmilled solid. During the calcination after the mixing with the additive, the copper is preferably replaced at least partly by the additional metal. The finished catalyst preferably comprises at least 70%, particularly preferably at least 80%, in particular at least 90%, of a spinel phase.

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As elements of main group II or transition group II of the Periodic Table of the Elements, it is possible to use not only oxides and the elements in metallic form but also their salts. Examples are carbonates, hydroxides, carboxylates, halides and oxidic anions such as nitrites, nitrates, sulfides, sulfates, phosphites, phosphates, pyrophosphates, halites, halates and basic carbonates. Preference is given to carbonates, hydroxides, carboxylates, nitrites, nitrates, sulfates, phosphates and basic carbonates, particularly preferably carbonates, hydroxides, basic carbonates and nitrates. The additional metal is particularly preferably in the oxidation state +2. Preference is given to using Zn, Mg, Ca, Sr and/or Ba, in particular Zn and/or Mg.

The preparation of the starting oxide of the composition CuAl₂O₄, preferably in the form of a spinel, is known from, for example, Z. Phys. Chem., 141 (1984), pages 101 to 103. Preference is given to impregnating an Al₂O₃ support with a solution of an appropriate salt. The anion is then preferably decomposed thermally to form the oxide. It is also possible to mix the salt with the aluminum compound (for example in suspension with subsequent spray drying), compact it and then bring it into the desired shape, followed by calcination.

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The catalyst preferably comprises from 0.1 to 30% by weight of CuO, from 0.1 to 40% by weight of the further metal oxide, in particular ZnO, and from 50 to 80% by weight of Al₂O₃.

The catalyst is particularly preferably made up of about 8% by weight of CuO, 30% by weight of ZnO and 62% by weight of Al_2O_3 . Apart from the spinel, preferably small amounts of CuO and further metal oxide are also present. Preferably, not more than 3.5% by weight of CuO and not more than 10% by weight of ZnO are present.

Suitable catalysts are also described in DE-A-43 01 469 and EP-A-0 687 499. Further examples of preparations of catalysts which can be used according to the present invention may be taken from the documents cited.

The catalyst preferably has a BET surface area of from 1 to 350 m^2/g . The porosity is preferably in the range from 0.01 to 0.8 l/g.

In the reactor of the present invention, the catalyst is preferably used in the star extrudate form described as a fixed bed. The thickness of the fixed bed is preferably from 2 to 50 cm, particularly preferably from 5 to 10 cm. The residence time over the catalyst for the decomposition of N_2O is preferably less than 0.1 s.

The use of the catalyst directly in the reactor for the catalytic oxidation of ammonia leads to complete degradation of N₂O, with nitrogen oxides being formed. The nitrogen oxides formed in the oxidation of ammonia are not degraded over this catalyst. The catalyst has a high activity. As a result of the low height of the catalyst bed and the preferred star extrudate shape of the catalyst, only a small pressure drop occurs in the reactor. No additional heating or cooling is required for the removal of N₂O. Since the reactors are built for accommodating catalyst gauzes, rebuilding of a nitric acid plant is generally not necessary.

The invention is illustrated by the examples below.

Example 1

In a laboratory apparatus, ammonia in an ammonia/air mixture having a concentration of 12.5% by volume of ammonia and 87.5% by volume of air was reacted over a Pt/Rh gauze at 913°C and a throughput of 37 g/h of ammonia per cm³ of gauze area to form nitrogen monoxide. Immediately downstream of the platinum gauze, there was a 10 cm high bed of the above-described preferred catalyst consisting of 8% by weight of CuO, 30% by weight of ZnO and 62% by weight of Al₂O₃; the reaction gas flowed through this bed at 750°C with a residence time of 0.03 s. Upstream of the catalyst bed, the N₂O concentration was 685 ppm, while only 108 ppm were present downstream of the catalyst bed. The content of nitrogen monoxide remained unchanged before and after the catalyst bed.

Example 2

In an atmospheric pressure reactor for the catalytic oxidation of ammonia to nitrogen oxides, which was equipped with a platinum/rhodium gauze as catalyst and a palladium gauze for noble metal recovery and a heat exchanger, a fixed catalyst bed having a thickness of 9 cm was installed between the recovery gauze and the heat exchanger. The catalyst bed comprised the above-described preferred catalyst consisting of 8% by weight of CuO, 30% by weight of ZnO and 62% by weight of Al₂O₃. The catalyst was used as star extrudates having a diameter of 4 mm. The reactor was supplied with a mixture of 12.1% by volume of ammonia and 87.9% by volume of air; the gas throughput corresponded to 15,000 kg/day of HNO₃ per m³ of noble metal gauze. The temperature in the reactor was 860°C. The residence time over the N₂O decomposition catalyst described was less than 0.05 s. The concentration of N₂O at the outlet of the reactor was 120 ppm.

In a comparative measurement, the additional catalyst bed for the decomposition of N_2O was omitted. An N_2O concentration at the reactor outlet of 606 ppm was obtained.

20 The measured NO_x concentration was in both cases 11.28% by volume.

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We claim:

- A reactor for the catalytic oxidation of ammonia to nitrogen oxides, which
 contains a noble metal gauze catalyst and a heat exchanger in that order in
 the direction of flow and has a catalyst for the decomposition of N₂O
 located between the noble metal gauze catalyst and the heat exchanger.
- A reactor as claimed in claim 1, wherein a noble metal recovery gauze is located between the noble metal gauze catalyst and the catalyst for the decomposition of N₂O.
 - A reactor as claimed in claim 1 or 2, wherein the catalyst for the decomposition of N₂O is installed as a fixed bed having a height of from 2 to 50 cm.
 - 4. A reactor as claimed in any of claims 1 to 3, wherein the catalyst for the decomposition of N₂O can be prepared by combining CuAl₂O₄ with tin, lead and/or an element of main group II or transition group II of the Periodic Table of the Elements as oxide or salt or in elemental form and subsequently calcining the mixture at from 300 to 1300°C and a pressure in the range from 0.1 to 200 bar.
- An apparatus for preparing nitric acid from ammonia, comprising in this
 order
 - a reactor as claimed in any of claims 1 to 4,
 - an absorption unit for the absorption of nitrogen oxides in an aqueous medium and, if desired,
 - a reduction unit for the selective catalytic reduction of nitrogen oxides.
 - 6. A process for the catalytic decomposition of N₂O in a gas mixture obtained in the preparation of nitric acid by catalytic oxidation of ammonia, where the N₂O is decomposed catalytically over a catalyst for the decomposition of N₂O, wherein the hot gas mixture obtained from the catalytic oxidation of ammonia is brought into contact with the catalyst for the decomposition of N₂O prior to subsequent cooling.

- A process as claimed in claim 6, wherein the residence time over the catalyst for the decomposition of N₂O is less than 0.1 s.
- 5 8. A process as claimed in claim 6 or 7, wherein the decomposition of N₂O is carried out at from 600 to 950°C and/or at a pressure in the range from 1 to 15 bar.
- 9. A process as claimed in any of claims 6 to 8, wherein the catalyst for the decomposition of N_2O can be prepared by combining $CuAl_2O_4$ with tin, lead and/or an element of main group II or transition group II of the Periodic Table of the Elements as oxide or salt or in elemental form and subsequently calcining the mixture at from 300 to 1300°C and a pressure in the range from 0.1 to 200 bar.
 - The use of a catalyst as defined in claim 9 for the decomposition of N₂O in a reactor for the catalytic oxidation of ammonia.

Abstract

- 5 In a reactor for the catalytic oxidation of ammonia to form nitrogen oxides which has a noble metal gauze catalyst and a heat exchanger in that order in the direction of flow, a catalyst for the decomposition of N₂O is located between the noble metal gauze catalyst and the heat exchanger.
- In a process for the catalytic decomposition of N₂O in a gas mixture obtained in the preparation of nitric acid by catalytic oxidation of ammonia, where the N₂O is decomposed catalytically over a catalyst for the decomposition of N₂O, the hot gas mixture obtained from the catalytic oxidation of ammonia is brought into contact with the catalyst for the decomposition of N₂O prior to subsequent cooling.

Declaration, Power of Attorney

Page 1 of 4

0050/048985

We (I), the undersigned inventor(s), hereby declare(s) that:

My residence, post office address and citizenship are as stated below next to my name,

We (I) believe that we are (I am) the original, first, and joint (sole) inventor(s) of the subject matter which is claimed and for which a patent is sought on the invention entitled

Catalytic decomposition of N_2O

the specification of which

[X]	is attached hereto.	
[]	was filed on	as
	Application Serial No.	
	and amended on	-
[x] was filed as PCT international application	
	Number <u>PCT/EP99/02544</u>	
	on 15.04.1999	
	and was amended under PCT Article 19	
	on	_ (if applicable).

We (I) hereby state that we (I) have reviewed and understand the contents of the above-identified specification, including the claims, as amended by any amendment referred to above.

We (I) acknowledge the duty to disclose information known to be material to the patentability of this application as defined in Section 1.56 of Title 37 Code of Federal Regulations.

We (I) hereby claim foreign priority benefits under 35 U.S.C. § 119(a)–(d) or § 365(b) of any foreign application(s) for patent or inventor's certificate, or § 365(a) of any PCT International application which designated at least one country other than the United States, listed below and have also identified below, by checking the box, any foreign application for patent or inventor's certificate, or PCT International application having a filing date before that of the application on which priority is claimed. Prior Foreign Application(s)

Application No.	Country	Day/Month/Year	Priority Claimed	
19819882.5	Germany	27 April 1998	[x] Yes [] N	Мo

Status (pending, patented,

We (I) hereby claim the benefit under Title 35, application(s) listed below.	, United States Codes, § 119(e) of any United States prov	isiona
(Application Number)	(Filing Date)	
(Application Number)	(Filing Date)	

We (I) hereby claim the benefit under 35 U.S.C. § 120 of any United States application(s), or § 365(c) of any PCT International application designating the United States, listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States or PCT International application in the manner provided by the first paragraph of 35 U.S.C. § 112, I acknowledge the duty to disclose information which is material to patentability as defined in 37 CFR § 1.56 which became available between the filing date of the prior application and the national or PCT International filing date of this application.

Application Serial No.	Filing Date	abandoned)

And we (1) hereby appoint Messrs. HERBERT. B. KEIL, Registration Number 18,967; and RUSSEL E. WEINKAUF, Registration Number 18,495; the address of both being Messrs. Keil & Weinkauf, 1101 Connecticut Ave., N.W., Washington, D.C. 20036 (telephone 202–659–0100), our attorneys, with full power of substitution and revocation, to prosecute this application, to make alterations and amendments therein, to sign the drawings, to receive the patent, and to transact all business in the Patent Office connected therewith.

We (I) declare that all statements made herein of our (my) own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issuing thereon.

Volker Schumacher NAME OF INVENTOR

Signature of Inventor

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